# **PCT**

# WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



## INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 6:
C01B 7/24, 7/20
A1
(11) International Publication Number: WO 98/27005
(43) International Publication Date: 25 June 1998 (25.06.98)
(21) International Application Number: PCT/GB97/03347 (81) Designated States: AL, AM, AT, AU, AZ, BA, BB, BG, BR,

(30) Priority Data:

9626329.8

(22) International Filing Date:

19 December 1996 (19.12.96) GB

16 December 1997 (16.12.97)

(71) Applicant (for all designated States except US): BRITISH NUCLEAR FUELS PLC [GB/GB]: Springfields, Salwick, Preston, Lancashire PR4 0XJ (GB).

(72) Inventors; and

- (75) Inventors/Applicants (for US only): WAKEFIELD, Leigh, Edward [GB/GB]; British Nuclear Fuels plc, Springfields, Salwick, Preston, Lancashire PR4 0XJ (GB). WHYTE, Barnaby, John [GB/GB]; British Nuclear Fuels plc, Springfield, Salwick, Preston, Lancashire PR4 0XJ (GB).
- (74) Agent: PAWLYN, Anthony, Neil; Urquhart-Dykes & Lord, Tower House, Merrion Way, Leeds LS2 8PA (GB).

B1) Designated States: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE, GH, HU, ID, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG).

#### Published

With international search report.

Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.

- (54) Title: DEVICE AND METHOD FOR THE STORAGE, TRANSPORTATION AND PRODUCTION OF ACTIVE FLUORINE
- (57) Abstract

An improved source of active fluorine is provided in the form of storing and transporting the active fluorine as alkali metal fluorohalates. The method generates the active fluorine at the point of use by thermally decomposing the source. The transportable form is safer to transport than previous active fluorine sources. Further methods involving the use of active fluorine produced in this way are also provided.

# FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

ΛL	Albania	ES	Spain	LS	Lesotho	SI	Slovenia
AM	Armenia	F1	Finland	LT	Lithuania	SK	Slovakia
AΤ	Austria	FR	France	LU	Luxembourg	SN	Senegal
AU	Australia	GA	Gahon	LV	Latvia	SZ	Swaziland
ΛZ	Azerbaijan	GB	United Kingdom	MC	Monaco	TD	Chad
BA	Bosnia and Herzegovina	GE	Georgia	MD	Republic of Moldova	TG	Togo
BB	Barbados	GH	Ghana	MG	Madagascar	ТJ	Tajikistan
BE	Belgium	GN	Guinea	MK	The former Yugoslav	TM	Turkmenistan
BF	Burkina Faso	GR	Greece		Republic of Macedonia	TR	Turkey
BG	Bulgaria	HU	Hungary	ML.	Mali	TT	Trinidad and Tobago
13,3	Benin	Æ	Ireland	MN	Mongolia	UA	Ukraine
BR	Brazil	H	Israel	MR	Mauritania	UG	Uganda
BY	Belains	IS	lcelaud	MW	Malawi	us	United States of America
CA	Canada	IT	italy	MX	Mexico	UZ	Uzbekistan
CF	Central African Republic	JР	Japan	NE	Niger	VN	Viet Nam
CG	Congo	KE	Kenya	NL	Netherlands	YU	Yugoslavia
CH	Switzerland	KG	Kyrgyzstan	NO	Norway	zw	Zimbabwe
CI	Côte d'Ivoire	KP	Democratic People's	NZ	New Zenland		
CM	Cameroon		Republic of Korea	PL	Poland		
CN	China	KR	Republic of Korea	PT	Portugal		
CU	Cuba	KZ	Kazakstan	RO	Romania		
CZ	Czech Republic	LC	Saint Lucia	RU	Russian Federation		
DE	Germany	LI	Liechtenstein	SD	Sudau		
DK	Denmark	LK	Sri Lanka	SE	Sweden		
EE	Estonia	LR	Liberia	SG	Singapore		

# WO 98/27005 PCT/GB97/03347

DEVICE AND METHOD FOR THE STORAGE. TRANSPORTATION AND PRODUCTION OF ACTIVE FLUORINE

This invention relates to the transportation, storage and production of active fluorine from safer sources and forms and in particular, but not exclusively to chlorine based active fluorine.

Active fluorine sources have many uses in industry. These include its use as a specialised fluorinating agent and rocket fuel as well as a high performance cleaning agent due to its extremely reactive nature. They can also be used to enhance the anti-corrosion properties of metals.

In semi-conductor manufacture they are used for etching silicon and as a cleaning agent. Semi-conductor and liquid crystal manufacture give rise to silicon and silicon oxide which adheres or deposits on the manufactured units. This material is effectively removed by active fluorine.

However, conventionally used sources of active fluorine in their form ready for use are highly dangerous substances and consequently there are exceedingly strict transportation regulations. The dangers with active fluorine sources arise from the fact that at room temperature and pressure they tend to be extremely reactive. Organic substances for example will react immediately with many active fluorine sources and ignite. Many active fluorine sources also react violently with water giving an emission of toxic fumes. Additionally many active fluorine sources react with any moisture in air and on contact with body tissue produce substances which are both toxic and corrosive to the body. Active fluorine sources are not only a toxic hazard but can also be a fire hazard.

Presently active fluorine sources are stored and transported in relatively small pressurised cylinders.

The present invention seeks to overcome the above-mentioned problems by providing a device and method for the provision of an active fluorine gas source at the point of use which is convenient and safer to store and transport.

S STAR I I

In accordance with one aspect of the present invention there is provided a device for producing active fluorine comprising :-

- i) a transportable vessel containing a solid alkali metal fluorohalate, preferably tetrafluorohalate,
- ii) means for heating the solid alkali metal fluorohalate, and
- iii) an outlet from the vessel from which the active fluorine gas produced on heating exits the vessel.

The alkali metal fluorohalate may be  $MXF_{(n+1)}$ , where M is K, Rb or Cs, X is Cl, Br, I and n is 1, 3, 5 or 7. A mixture of alkali metal fluorohalates may be provided.

The active fluorine may be or may include F,  $F_2$ , ClF, ClF<sub>3</sub>, ClF<sub>5</sub>, IF<sub>5</sub>, IF<sub>7</sub>, BrF, BrF<sub>3</sub>, BrF<sub>5</sub> or mixtures thereof. The gas produced may include Cl<sub>2</sub>, Cl, Br<sub>2</sub>, Br, I, I<sub>2</sub> or mixtures thereof.

Active fluorine consists of highly energetic fluorine species, such as a fluorine radical.

The fluorohalate, and preferably tetrafluorohalate, may be bromate or iodate but is preferably chlorate.

Any alkali metal may be employed. Preferably the solid alkali metal tetrafluorochlorate is  $KClF_4$ , or alternatively it may be  $RbClF_4$ , or  $CsClF_4$ , or any other suitable alkali metal which may provide a solid suitable for transportation.

The alkali metal fluorohalate, such as tetrafluorohalate is preferably provided in pellet form within the transportable vessel. The pellets may be provided on support layers.

The transportable vessel is preferably a cylinder which is adapted to be able to contain the solid alkali metal fluorohalate. A nickel or monel lined cylinder may be used. A filter may be provided, most preferably at the outlet, to help retain the solids. A sintered filter may be preferred.

The alkali metal fluorohalate may be charged to a cartridge capable of insertion into the transportable vessel.

The heater is preferably an integral part of the vessel so that the gas may be produced easily at any point of use. The heater may be provided in the form of heat transfer fins made for example of suitable metal strips. These fins may be provided in the vessel and may extend radially outwards from the centre of

the vessel providing an effective heat source to the core of the vessel. Alternatively the heater may be provided separately and applied at the point of use, as required when the production of the active fluorine gas is needed.

An outlet for the vessel may preferably be an industry standard fitting or flange.

The device may be provided with a control system to regulate the temperature and/or pressure generated by heating.

The device may be recharged following use. The device may be recharged using material extracted from the waste stream following the use of the active fluorine, for instance from a scrubber such as a KF scrubber system which selectively absorbs the active fluorine in a re-releasable form.

According to a second aspect of the present invention there is provided a method of providing a source of active fluorine comprising:-

- i) generating a solid alkali metal fluorohalates, preferably tetrafluorohalate,
- ii) transporting and storing the source as a solid alkali metal fluorohalate, and
- iii) heating the fluorohalate at the point of use to generate active fluorine gas.

The active fluorine may comprise fluorine alone or in combination with any halogen but fluorine.

The active fluorine may be or may include F,  $F_2$ , ClF, ClF<sub>3</sub>, ClF<sub>5</sub>, IF<sub>5</sub>, IF<sub>7</sub>, BrF, BrF<sub>3</sub> or mixtures thereof. The gas produced may include Cl<sub>2</sub>, Cl, Br<sub>2</sub>, Br, I, I<sub>2</sub> or mixtures thereof.

Active fluorine consists of highly energetic fluorine species, such as a fluorine radical.

The fluorohalate, and preferably tetrafluorohalate, may be icdate or bromate, but is preferably the chlorate form.

Any alkali metal may be employed. The solid alkali metal tetrafluorochlorate to be produced is preferably KClF<sub>4</sub>, but it may alternatively be RbClF<sub>4</sub>, or CsClF<sub>4</sub>, or any other suitable alkali metal tetrafluorochlorate.

In order to produce active fluorine from the  $MClF_4$ , (where M equals alkali metal) contained within the vessel, the solid is

WO 98/27005

heated to a temperature at which the  $MClF_4$ , decomposes to form active fluorine gas.

The solid alkali metal tetrafluorochlorate is preferably heated to a temperature of at least 90°C or 100°C to cause the generation of active fluorine. Alternatively temperatures of at least 130°C, or at least 180°C may be used. Preferably a temperature of 90°C to 300°C or 325°C is used to decompose the alkali metal tetrafluorochlorate to form active fluorine gas, on demand.

Preferably the decomposition is carried out in a vacuum or in an inert gas stream or merely in the presence of active fluorine.

According to a third aspect of the invention we claim the use of alkali metal fluorohalate, preferably tetrafluorohalate, as a transportable and/or storable source of active fluorine.

The use of alkali metal tetrafluorochlorate as a source of active fluorine is preferred. This technique offers significant safety advantages as the inventory of free active fluorine is kept to a minimum.

According to a fourth aspect of the invention we provide the use of active fluorine as a cleaning agent and/or etching agent.

Preferably the active fluorine is produced using the device of the first aspect and/or the method of the second aspect.

Preferably the active fluorine is used for cleaning chemical vapour deposition equipment. The active fluorine may be used for removal of silicon and/or germanium and/or copper and/or compounds thereof, from semi-conductor manufacturing apparatus, liquid crystal manufacturing apparatus or associated accessories thereof.

The removal of silicon is particularly preferred. The active fluorine may be used for etching silicon, germanium, copper or compounds thereof.

The active fluorine may be used to clean other apparatus or equipment or locations, such as piping.

According to a fifth aspect of the invention we provide a method of cleaning and / or etching comprising producing active fluorine by the device of the first aspect of the invention and / or according to the method of the second aspect of the

invention and applying the active fluorine to a material to be cleaned or etched.

-5-

PCT/GB97/03347

According to a sixth aspect of the invention we provide a method of passivating metals by fluorination using active fluorine generated using the device of the first aspect of the invention and/or using the method of the second aspect of the invention.

The invention will now be described in detail, by way of example only, and with reference to the following figures in which :-

Figure 1 is an illustration of a storage unit according to the invention; and

Figure 2 is a schematic diagram illustrating the provision of the active fluorine in accordance with the present invention.

Solid KClF4, within the vessel may be transported and stored wherever the active fluorine is required with free active fluorine only being generated when there is a demand for such gas. The provision of the active fluorine in this way offers a number of benefits over the prior art metallic cylinders containing active fluorine or sources thereof. Firstly the KClF4, is a comparatively stable substance, when compared with other active fluorine sources, giving a higher level of safety during transport and storage. Its ease of loading into the transport means is also increased as cartridges incorporating the material, a solid, can be more readily provided. effectiveness as a source of cleaning gas is also higher than prior art sources such as fluorine ions. Equally the materials can be employed at room temperatures as a plasma is no longer needed. Significantly the active fluorine source is also free of carbon materials avoiding CFC formation and release.

Figure 1 illustrates a cylinder according to an embodiment of the invention in which the cylinder 1 is provided at each end with a valve 2a, 2b. The solid KClF4, 10, is stored within the cylinder 1 and a sintered filter 3 is provided between the solid and outlet valve 2a. Heating elements 4 are wrapped around or placed against the cylinder 1 and provided with power via leads 5. A thermocouple is provided to monitor the heating. Encasing the assembly is a layer of thermal insulation 6 which itself is contained within a structural safety jacket 7.

The present invention is exemplified by the generation of active fluorine from a device as illustrated in Figure 2.

In order to obtain the active fluorine gas at the point of use, Figure 2, the cylinder 1 is connected to the location of use 100 via a pipe 102 incorporating a control valve 104. Electrical connections are attached to the leads 5 and a power supply 106 attached.

Under the control of a heater switch power 104 is introduced to the heating elements 4. The  $KClF_4$  as a consequence is heated to a temperature of up to 300°C, and preferably 230°C, by the integral heating elements 4. The heating is monitored by the use of a thermocouple. At this temperature the solid releases some gas and the gas is released into the pipe. The filter in the cylinder restrains any solid. The pipe 102 directs the gas to the location 100 required.

The heater switch 104 can be controlled by a signal from a pressure monitor 108 connected to the gas outlet pipe 102 to maintain steady production.

Different decomposition temperatures are employed for other alkali metal tetrafluorochlorates; higher temperatures being employed for Caesium for instance.

# Embodiment 1

When the cartridge charged with the granular agent (equivalent of 100g of  $KClF_4$ ) carrying  $KClF_4$  on the Ni carrier using the unit of this invention as shown in Figure 1 is gradually heated by the heater elements 4 from outside with its outlet valve 2a closed, the rise of pressure inside the cartridge was observed, and a level of 1 kgf/cm² was observed at  $220\,^{\circ}$ C. When the outlet valve 2a was opened to allow flowrate of 2ml/min whilst keeping the temperature at  $240\,^{\circ}$ C, generation of gas was confirmed by a float system flow meter. The generation of gas was continued in that state. However, when the heater was switched off, a drop in the flow rate was observed as the temperature decreased.

For information, the cartridge to be charged with the agent should preferably be made of heat resistant and anti-corrosive materials such as Monel and Ni, but stainless steel such as SUS 316L can also be used.

# Embodiment 2

Gas was generated for a while by the use of the same unit as that in Embodiment 1 with the temperature kept at 240°C. Thereafter the heater was switched off and disconnected, and the air feed cartridge was cooled down from the outside. As a result, the outlet pressure became the atmospheric pressure at 220°C, and generation of gas stopped.

When the cartridge was heated again, gas was generated at 240°C, and the generation pressure increased as the temperature rose.

When the composition on the internal surface of cartridge was analyzed after sufficiently replacing the gas inside the cartridge with  $N_2$ , after the end of gas generation and returning temperature to the room temperature, the material found there was only fluorine, except the constituents of stainless steel SUS 316L which was the cartridge material. Chloride could hardly be detected.

## Embodiment 3

When the cartridge, charged this time with 50g of granular  $KClF_4$ , was gradually heated with the heater from outside, while a high purity  $N_2$  was flowed at the rate of 20 ml/minute from its bottom side 2b using the same unit as that in Embodiment 1, generation of active fluorine was observed at 120°C.

When the temperature was raised up to 180°C, a rise in concentration of relative active fluorine could be observed as the temperature rose. The relative concentration of active fluorine was measured by detecting hydrogen fluoride using a detection tube.

Examples of the use of active fluorine produced in this way and its benefits compared with prior art sources are now

exemplified, firstly with reference to the fluorination of steel and subsequently with reference to iron, nickel, aluminium, magnesium and finally chromium.

## Embodiment 4

EP 1/2 inch piping made of austenite stainless steel SUS 316L, was treated with diluted fluoric acid to removing the natural oxidized film, washed with water and dried for use as a test piece.

After moisture was removed by baking at 150°C, again while flowing a high purity  $N_2$  stream first, the gas supplied from the active fluorine generating unit was flowed at room temperature to performing fluorination treatment. For stabilizing the film formed on the internal face of SUS 316L at this time, heat treatment was made at 350°C while flowing a high purity  $N_2$  for forming a stable passive state film. The film compound was found by XDF thin film formation method to be mainly FeF $_2$ , and it was confirmed that the metallic compound (percentage) of SUS 316L remained almost unchanged.

In addition, observation of the surface after sealing with water containing 100 wtppm hydrogen bromide applied to the EP 1/2 inch column, made similarly of SUS 316L, at 50°C for 1 week gave no change on visual inspection, nor when observed by the XDF thin film method.

#### Comparison example 1

Fluorination of stainless steel SUS 316L.

EP 1/2 inch pipe made of austenitic stainless steel SUS 316L, was treated with the hydrofluoric acid to remove the natural oxidized film, washed with water and dried for use as a test piece.

After removing the moisture by baking the test piece at 250°C while flowing a high purity  $N_2$  stream first, the high purity hydrofluoric acid was flowed at 220°C to perform the fluorination treatment. At this time, to stabilize the film formed on the internal face of the SUS 316L piping, heat treatment at 350°C while flowing a high purity  $N_2$  stream was applied. The film

-9-

composition was mainly the FeF<sub>2</sub>, which was measured by XDF thin film method, and it was confirmed that the metal composition (percentage) of SUS 316L remained almost unchanged.

Further, when the surface of an EP 1/2 inch column made of SU3 316L that was prepared similarly was observed after sealing with water containing 100 wtppm hydrogen bromide, at 50°C for one week there was no visible change nor any change of composition observed even with the XDF thin film method.

However, when the fluorination treatment was conducted at the same room temperature as that of Embodiment 1, a perfect  $FeF_2$  film was not formed, and fluorination was found to be insufficient.

#### Embodiment 5

Fluorination of iron, nickel, aluminium and magnesium

Tubes of iron, nickel, aluminium and magnesium of purity of 99.9% or higher, were respectively treated with diluted hydrofluoric acid (then nitric acid + ammonium fluoride for the aluminium) to remove the natural oxidized film, washed with water and dried for use as the test pieces.

After removing the water content by baking at 150°C, again while flowing a high purity  $N_2$  stream first, the gas supplied from the active fluorine generating unit was flowed at room temperature to perform the fluorination treatment. In order to stabilize the film formed on the internal face of each metallic tube at this time, a stable passive film was formed by heat treatment at 350°C, while flowing a high purity  $N_2$  stream. The film compositions were confirmed to be mainly  $FeF_2$ ,  $NiF_2$ ,  $AIF_3$  and  $MgF_3$  by XDF thin film method.

# Embodiment 6

Fluorination of chrome.

A chrome tube of purity of 99.97 or higher, was treated with diluted hydrofluoric acid to remove the natural oxidised film, washed with water and dried for use as the test piece.

After removing the moisture content by baking at 150°C after again initially flowing a high purity  $N_2$  stream, the gas supplied

together with the high purity  $N_2$  from the active hydrofluoric acid generating unit like that in Embodiment 3 was flowed at room temperature to perform the fluorination treatment. At this time, for stabilizing the film formed on the internal face of the chrome tube, a stable passive film was formed by heat treatment at 200°C while flowing a high purity  $N_2$  stream.

The film composition was confirmed to be mainly  ${\rm Cr} F_3$  by the XDF thin film method.

Because chrome produces volatile  $CrF_5$  and splashes under fluorination conditions of metal in Embodiment 5, a stable passive film can not be formed.

The use of active fluorine produced by the present invention is now illustrated in terms of the etching and cleaning of silicon, silicon oxide, silicon nitride and silicon carbide. Again comparison with fluorine produced from prior art sources is provided.

## Embodiments 7 and 8

Etching and cleaning of silicon.

One face of an n-type silicon wafer, which was masked with the PMMA-based resin, and cut out to the size of 5  $\times$  5 mm and whose surface, fat/grease/oil, oxidation and stains were removed, was used as a test piece.

The test piece just after its preparation was installed inside the plasma CVD unit, and was made to react by introducing the undiluted active fluorine at a pressure of 5 Torr at a the flow rate of 30 SCCM at room temperature with the high frequency power source switched off.

Table 1 shows the result of investigating the reduction in mass of silicon for a gas passing time of 60 and 100 minutes. The  $F^{-}$  in the gas name column indicates the active fluorine generated by this invention.

# Comparison Examples 2 to 6:

Etching and cleaning of silicon

Table 1 shows the result of reacting the fluorine, nitrogen trifluoride, carbon tetrafluoride, dicarbonhexafluoride, a mixed gas of 95% carbon tetrafluoride and 5% oxygen, and a mixed gas of 5% dicarbonhexfluoride and 5% oxygen for the gas passing time of 100 minutes under the same conditions and using the same test pieces as those of Embodiments 7 and 8.

Circuit Gas Name Gas passing Reduction Remarks Board time mass F. Embodiment 60 min 16.2 mg Embodiment Si 100 min 22.5 mg Comparison 100 min < 0.1 mgSi  $F_2$ example 2 NF, 100 min < 0.1 mgComparison Si example 3 CF. 100 min < 0.1 mgComparison Si example 4 CF. 95%+ O<sub>2</sub> 5% 100 min Si < 0.1 mgComparison example 5 C2F6 95%+ Comparison Si 100 min < 0.1 mq0, 5% example 6

Table 1

## Embodiments 9 to 11:

Etching and cleaning of silicon oxide, silicon nitride and silicon carbide.

Layers of silicon oxide, silicon nitride and silicon carbide, with a thickness of 10,000 A, were formed respectively on SUS 316L plates of 100 x 100 x lmm thickness for use as the test pieces.

The test piece was installed inside the plasma CVD unit and reacted by introducing undiluted active fluorine at a pressure of 5 Torr at a flow rate of 30 SCCM at a temperature of 100°C with the high frequency power source switched off.

Table 2 shows the result of the investigations in terms of the reduction in mass of silicon oxide, silicon nitride and silicon carbide observed for a gas passing time of 3 minutes.

# Comparison Examples 7 to 15

Etching and cleaning of silicon oxide, silicon nitride and silicon carbide

Table 2 shows the result of reacting fluorine, nitrogen trifluoride, a mixed gas of 95% carbon tetrafluoride and 5% oxygen, and a mixed gas of 5% dicarbon hexafluoride and 5% oxygen under the same conditions and for a gas passing time of 100 minutes as using the same test pieces as those of Embodiments 9 to 11.

Table 2

	Circuit Board	Gas Name	Gas Passing Time	Reduction mass	Remarks
Embodiment 9	Silicon oxide	F'	3 min	14.2 mg	The circuit board was partially exposed.
Comparison example 7	Silicon oxide	F <sub>2</sub>	100 min	0.3 mg	
Comparison example 8	Silicon oxide	NF <sub>3</sub>	100 min	<0.1 mg	
Comparison example 9	Silicon oxide	CF <sub>4</sub> 95%+ O <sub>2</sub> 5%	100 min	<0.1 <b>m</b> g	
Embodiment 10	Silicon nitride	F'	3 min	33.7 mg	The circuit board was almost entirely exposed.
Comparison example 10	Silicon nitride	F <sub>2</sub>	100 min	1.4 mg	
Comparison example 11	Silicon nitride	NF <sub>3</sub>	100 min	<0.1 mg	
Comparison example 12	Silicon nitride	CF, 95%+ O <sub>2</sub> 5%	100 min	<0.1 mg	
Embodiment	Silicon carbide	F'	3 min	რ.6 mg	
Comparison example 13	Silicon carbide	F <sub>2</sub>	100 min	0.1 mg	
Comparison example 14	Silicon carbide	NF <sub>3</sub>	100 min	<0.1 mg	
Comparison example 15	Silicon carbide	CF, 95% O, 5%	100 min	<0.1 mg	

-13-

# CLAIMS

- A device for producing active fluorine comprising :-
- i) a transportable vessel containing a solid alkali metal fluorohalate, preferably tetrafluorohalate,
- ii) means for heating the solid alkali metal fluorohalate, and
- iii) an outlet from the vessel from which the active fluorine gas produced on heating exits the vessel.
- A device according to claim 1 in which the alkali metal fluorohalate is  $MXF_{(n+1)}$ , where M is K, Rb or Cs, X is Cl, Br, I and n is 1, 3, 5 or 7.
- 3. A device according to claim 1 or claim 2 in which the fluorohalate is  $KClF_4$  or  $RbClF_4$  or  $CsClF_4$ .
- 4. A device according to any of claims 1 to 3 in which the transportable vessel is a cylinder which is adapted to be able to contain the solid alkali metal tetrafluorohalate.
- 5. A device according to any of claims 1 to 4 in which the heater is an integral part of the vessel.
- 6. A device according to any of claims 1 to 5 in which the device is recharged following use.
- providing a source of active fluorine method of 7. A comprising: -
- i) generating a solid alkali metal fluorohalates, preferably tetrafluorohalate,
- ii) transporting and storing the source as a solid alkali metal fluorohalate, and
- iii) heating the fluorohalate at the point of use to generate active fluorine gas.
- The method of claim 7 in which the active fluorine comprises fluorine alone or in combination with any halogen but fluorine,

for instance the fluorine may be or may include F,  $F_2$ ,  $ClF_3$ ,  $ClF_5$ ,  $IF_5$ ,  $BrF_3$  or mixtures thereof.

- 9. A method according to claim 7 or claim 8 in which the the  $MClF_4$ , (where M equals alkali metal) contained within the vessel is heated to a temperature at which the  $MClF_4$ , decomposes to form active fluorine gas.
- 10. A method according to claim 9 in which the solid alkali metal tetrafluorchlorate is heated to a temperature of at least 130°C, or at least 180°C.
- 11. A method according to any of claims 7 to 10 in which the decomposition is carried out in a vacuum or in an inert gas stream or in the presence of active fluorine.
- 12. The use of alkali metal fluorohalate, preferably tetrafluorohalate, as a transportable and/or storable source of active fluorine.
- 13. A method of cleaning and/or etching comprising producing active fluorine by a device according to any of claims 1 to 6 and/or according to the method of any of claims 7 to 11 and applying the active fluorine to a material to be cleaned or etched.
- 14. A method according to claim 13 in which the active fluorine is used for cleaning chemical vapour deposition equipment and/or for removal of silicon and/or germanium and/or copper and/or compounds thereof, from semi-conductor manufacturing apparatus, liquid crystal manufacturing apparatus or associated accessories thereof, and/or for cleaning piping.
- 15. A method according to claim 13 or claim 14 in which the active fluorine is used for etching silicon, germanium, copper or compounds thereof.

PCT/GB97/03347

-15-

16. A method of passivating metals by fluorination using active fluorine generated using the device of any of claims 1 to 6 and/or using the method of any of claims 7 to 11.

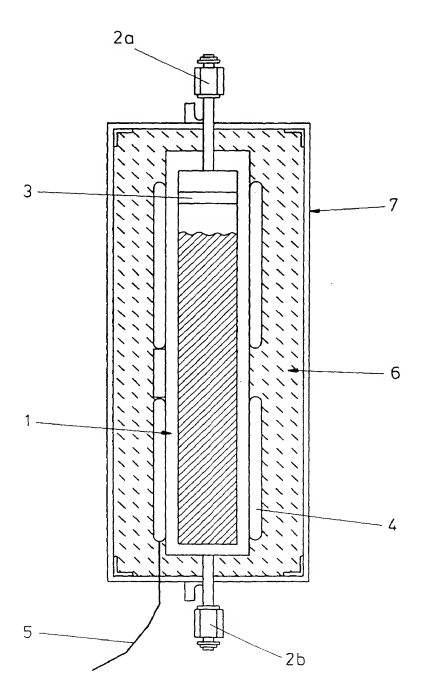


FIG. 1

SUBSTITUTE SHEET (RULE 26)

PCT/GB97/03347 WO 98/27005

-2/2-

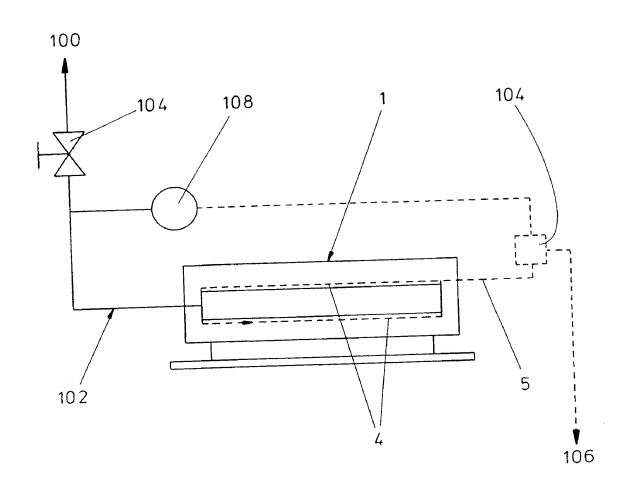


FIG. 2

## INTERNATIONAL SEARCH REPORT

Inte Jonal Application No PCT/GB 97/03347

A. CLASSIFICATION OF SUBJECT MATTER IPC 6 CO187/24 CO18 C01B7/20 According to International Patent Classification (IPC) or to both national classification and IPC **B. FIELDS SEARCHED** Minimum documentation searched (classification system followed by classification symbols) IPC 6 CO1B Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Category ' Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No Α CHEMICAL ABSTRACTS, vol. 85, no. 12, 1 20 September 1976 Columbus, Ohio, US; abstract no. 80474, "Solid reactants for FOGLE, C. E. ET AL: fluorine gas generators" XP002062746 see abstract U. S. NTIS, AD REP. (1976), AD-A022099, 112 PP. AVAIL.: NTIS FROM: GOV. REP. ANNOUNCE. INDEX (U. S.) 1976, 76(10), 211 CODEN: XADRCH, 1976, US 3 361 543 A (TSIGDINOS GEORGE A ET AL) Α 1 2 January 1968 see claim 1 -/--Further documents are listed in the continuation of box C Patent family members are listed in annex Х Special categories of cited documents : \*T\* later document published after the international filing date or priority date and not in conflict with the application but \*A\* document defining the general state of the art which is not cited to understand the principle or theory underlying the considered to be of particular relevance \*E\* earlier document but published on or after the international \*X\* document of particular relevance; the claimed invention filing date cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such docu-\*O\* document referring to an oral disclosure, use, exhibition or ments, such combination being obvious to a person skilled in the art. \*P\* document published prior to the international filing date but later than the priority date claimed \*&\* document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report 21 April 1998 13. 05 199A Name and mailing address of the ISA Authorized officer European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo ni, Clement, J-P Fax: (+31-70) 340-3016

Form PCT/ISA/210 (second sheet) (July 1992)

1

# INTERNATIONAL SEARCH REPORT

Inter onal Application No
PCT/GB 97/03347

		PCT/GB 97/03347	
Continuategory °	ation) DOCUMENTS CONSIDERED TO BE RELEVANT  Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.	
icego. y	oration of account, with management, where appropriate, or the factoring passages		
	US 4 284 617 A (BOWEN RICHARD E ET AL) 18 August 1981 see claims 11,12	1	
1	US 3 143 391 A (HURLEY THOMAS J ET AL) 4 August 1964 see claim 3	1	
	US 3 989 808 A (ASPREY LARNED B) 2 November 1976 see claim 1	1	
	·		

1

Form PCT/ISA/210 (continuation of second sheel) (July 1992)

# INTERNATIONAL SEARCH REPORT

•

Information on patent family members

Inte onal Application No PCT/GB 97/03347

Patent document cited in search report	Publication date	Patent family member(s)	Publication date	
US 3361543 A	02-01-68	NONE		
US 4284617 A	18-08-81	NONE		
US 3143391 A	04-08-64	NONE		
US 3989808 A	02-11-76	NONE		

Form PCT/ISA/210 (patent family annex) (July 1992)